

University of Groningen

SpotXplore

Westenberg, Michel A.; Roerdink, Jos B. T. M.; Kuipers, Oscar P.; van Hijum, Sacha A. F. T.

Published in:
Bioinformatics

DOI:
[10.1093/bioinformatics/btq535](https://doi.org/10.1093/bioinformatics/btq535)

IMPORTANT NOTE: You are advised to consult the publisher's version (publisher's PDF) if you wish to cite from it. Please check the document version below.

Document Version
Publisher's PDF, also known as Version of record

Publication date:
2010

[Link to publication in University of Groningen/UMCG research database](#)

Citation for published version (APA):

Westenberg, M. A., Roerdink, J. B. T. M., Kuipers, O. P., & van Hijum, S. A. F. T. (2010). SpotXplore: A Cytoscape plugin for visual exploration of hotspot expression in gene regulatory networks. *Bioinformatics*, 26(22), 2922-2923. <https://doi.org/10.1093/bioinformatics/btq535>

Copyright

Other than for strictly personal use, it is not permitted to download or to forward/distribute the text or part of it without the consent of the author(s) and/or copyright holder(s), unless the work is under an open content license (like Creative Commons).

The publication may also be distributed here under the terms of Article 25fa of the Dutch Copyright Act, indicated by the "Taverne" license. More information can be found on the University of Groningen website: <https://www.rug.nl/library/open-access/self-archiving-pure/taverne-amendment>.

Take-down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

Downloaded from the University of Groningen/UMCG research database (Pure): <http://www.rug.nl/research/portal>. For technical reasons the number of authors shown on this cover page is limited to 10 maximum.

SpotXplore: a Cytoscape plugin for visual exploration of hotspot expression in gene regulatory networks

Michel A. Westenberg^{1,*}, Jos B.T.M. Roerdink², Oscar P. Kuipers³
and Sacha A.F.T. van Hijum^{4,5,6,7}

¹Department of Mathematics and Computer Science, Eindhoven University of Technology, Eindhoven, ²Johann Bernoulli Institute for Mathematics and Computer Science, University of Groningen, Groningen, ³Department of Genetics, University of Groningen, Haren, ⁴Center for Molecular and Biomolecular Informatics, Radboud University Nijmegen Medical Centre, Nijmegen, ⁵Kluyver Centre for Genomics of Industrial Fermentation, NIZO food research, Ede, ⁶TI Food and Nutrition, Wageningen and ⁷Netherlands Bioinformatics Centre, Nijmegen, The Netherlands

Associate Editor: Trey Ideker

ABSTRACT

Summary: SpotXplore is a plugin for Cytoscape for extraction and visualization of differentially expressed subnetworks (hotspots) from gene networks. The hotspot-based visualization approach enables interactive exploration of regulatory interactions in differentially expressed gene sets, and it allows a researcher to explore gene expression in direct relation to the affected cellular gene network. The hotspots provide a view beyond the commonly used metabolic pathways and gene ontologies.

Availability: <http://www.win.tue.nl/~mwestenb/spotxplore/>

Contact: m.a.westenberg@tue.nl

Supplementary information: Example data and user manual are available at SpotXplore's web site.

Received on June 4, 2010; revised on September 8, 2010; accepted on September 15, 2010

1 INTRODUCTION

Interpretation of gene expression from time series experiments or multiple perturbation studies is a challenging task. Insight can be obtained by analyzing the data in the context of molecular interaction networks, such as gene regulatory networks or metabolic pathways, which requires visualization in addition to statistical analysis tools. Many tools for visualizing the various types of biological networks have been developed (Suderman and Hallett, 2007). Nevertheless, there is a need for tools that incorporate dynamics of gene expression in the networks by offering both data analysis and visualization. Only then, a researcher does not have to switch between various applications to obtain a comprehensive understanding of the biological processes affected in the experiments.

The generic biological network visualization tool Cytoscape (Shannon *et al.*, 2003) provides an excellent environment to implement combined analysis and visualization methods. In this article, we present SpotXplore, a plugin for analysis and visual exploration of differentially expressed subnetworks (hotspots). SpotXplore provides two methods to detect such hotspots, and allows the user to visualize the hotspots in the context of the whole network to explore regulatory interactions between differentially

expressed gene sets. Our plugin also provides a method for visualization of time series data, because Cytoscape offers only basic support by coloring nodes (genes) based on a single time point. However, to detect temporal patterns in gene expression, it is necessary to consider all time points, which requires simultaneous visualization of multiple expression values. For this purpose, we re-implemented and adapted the visualization approach of our stand-alone tool GENeVis (Westenberg *et al.*, 2007), which allows embedded visualization of all time points in the nodes. To our knowledge, only the VistaClara plugin (Kincaid *et al.*, 2008) offers a similar visualization by drawing time points *under* the nodes. Our approach is different, because it also encodes statistical properties of the time points in the visualization, which is important for data analysis.

SpotXplore's main contribution is the hotspot-based visualization approach, which enables interactive exploration of regulatory interactions in differentially expressed gene sets, allowing a researcher to explore gene expression in direct relation to the affected cellular gene network.

2 METHODS AND IMPLEMENTATION

SpotXplore is a plugin for Cytoscape, which has been tested in versions 2.6.3 and 2.7.0 on multiple platforms (Windows, Linux, MacOS). It requires a gene network and gene expression data already loaded in Cytoscape. Gene networks can be obtained from databases, such as DBTBS (Makita *et al.*, 2004), or by constructing a co-regulation network from the expression data. The user manual, available as Supplementary Material, provides details about the required and optional node and edge attributes.

2.1 Gene expression visualization

SpotXplore assumes that gene expressions are given as log-transformed ratios between a test condition and a reference condition. Each node (representing a gene) can be colored according to the expression ratio. Additionally, all gene expression ratios for a certain gene, i.e. all time points or all conditions, can be shown simultaneously as small glyphs embedded in the rectangular box representing the gene. Each glyph is a small rectangle, colored according to expression ratio and scaled in height according to confidence value (e.g. a *P*-value) (Westenberg *et al.*, 2007). This graphical representation is inspired by perception research, from which it is known that color and size are perceptually easy to separate and interpret independently.

*To whom correspondence should be addressed.

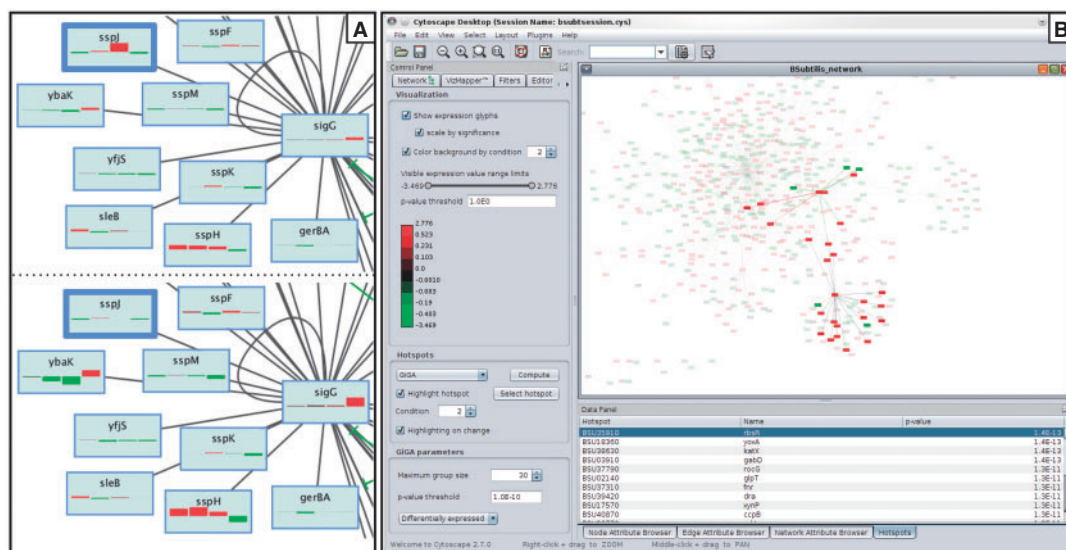


Fig. 1. SpotXplore applied to a *Bacillus subtilis* gene network and a four time point microarray experiment (Supplementary Materials). (A) Part of the SigG regulon. Expression ratio and *P*-values of four time points are displayed in the nodes. Time point boxes are colored according to expression ratio (red–green). Top: box height scaled linearly by expression ratio. Bottom: box height scaled logarithmically by *P*-value. Scaling by expression ratio may give the impression that the gene *sspJ* (highlighted by a wide border) is strongly upregulated at time point 3. However, the corresponding *P*-value is 1.0. Scaling by significance value suppresses these insignificant data points (see bottom). Conversely, it enhances data points that do show significant differential expression. (B) Screenshot of SpotXplore. Left panel: visual appearance controls, color legend and hotspot detection algorithm parameters. Bottom right panel: table of detected hotspots. The hotspot *rrsR* at time point 2 is selected and its members highlighted (top right panel). The remaining nodes (genes that are not differentially expressed) and edges are drawn translucent.

Scaling by significance gives reliable time points more visual emphasis and suppresses noisy expression ratios (Fig. 1A).

2.2 Hotspot detection

Hotspots are network components that are differentially expressed as a function of experimental parameters. We provide two algorithms to detect hotspots: Graph-based iterative Group Analysis (Breitling *et al.*, 2005) and a two-pass algorithm that requires both expression ratios and expression levels for its computation (Luscombe *et al.*, 2004). Each algorithm computes hotspots per condition or time point. SpotXplore runs the selected algorithm for all conditions, and stores the detected hotspots for later exploration. Additionally, hotspots can be defined by importing a set of selected nodes from Cytoscape. This allows quick manual construction of a set of hotspots, and also provides a way to use our visualization approach for hotspots detected by other Cytoscape plugins.

2.3 Hotspot visualization

SpotXplore visualizes hotspots in the context of the whole network (Fig. 1B). The conditions (or time points) can be browsed and an overview of the network activity is provided by highlighting all hotspots detected in a certain condition. An individual hotspot can be highlighted by selecting it, and can be converted into a Cytoscape selection set for further analysis.

3 CONCLUSION

SpotXplore is a plugin for the widely adopted generic Cytoscape network visualization software. It allows to interactively explore

regulatory interactions in differentially expressed gene sets, occurring in collections of gene expression data (e.g. time series), obtained by microarrays or RNA-seq techniques. The approach both facilitates and significantly speeds up data analysis in comparison to a traditional gene-by-gene analysis approach of the most differentially expressed genes and manual mining of gene regulatory interaction information available at public repositories.

Conflict of Interest: none declared.

REFERENCES

- Breitling, R. *et al.* (2005) Graph-based iterative group analysis enhances microarray interpretation. *BMC Bioinformatics*, **5**, 100.
- Kincaid, R. *et al.* (2008) VistaClara: an expression browser plug-in for Cytoscape. *Bioinformatics*, **24**, 2112–2114.
- Luscombe, N.M. *et al.* (2004) Genomic analysis of regulatory network dynamics reveals large topological changes. *Lett. Nat.*, **431**, 308–312.
- Makita, Y. *et al.* (2004) DBTBS: database of transcriptional regulation in *Bacillus subtilis* and its contribution to comparative genomics. *Nucleic Acids Res.*, **32**, D75–D77.
- Shannon, P. *et al.* (2003) Cytoscape: a software environment for integrated models of biomolecular interaction networks. *Genome Res.*, **13**, 2498–2504.
- Suderman, M. and Hallett, M. (2007) Tools for visually exploring biological networks. *Bioinformatics*, **23**, 2651–2659.
- Westenberg, M.A. *et al.* (2007) Interactive visualization of gene regulatory networks with associated gene expression time series data. In Linsen, L. *et al.* (eds), *Visualization in Medicine Life Sciences, Visualization and Mathematics*. Springer, Berlin, Germany, pp. 293–312.